

SEQUENCE LISTING

<110> Gray et al.

<120> MACROPHAGE DERIVED CHEMOKINE (MDC), MDC ANALOGS, MDC INHIBITOR SUBSTANCES, AND USES THEREOF

<130> 27866/34810

<140> 09/509,165

<141> 2000-06-12

<150> 09/067,447

<151> 1998-04-28

<150> 08/939,107

<151> 1997-09-26

<150> 08/660,542

<151> 1996-06-07

<150> 08/558,658

<151> 1995-11-16

<150> 08/479,620

<151> 1995-06-07

<160> 46

<170> PatentIn Ver. 2.0

<210> 1

<211> 2923

<212> DNA

<213> Homo sapiens - human MDC cDNA

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<221> CDS

<222> (20)..(298)

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<221> mat_peptide

<222> (92)..(298)

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-10

- 1

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                                    -15
Val Ala Leu Gln Ala Thr Glu Ala Gly Pro/ Tyr Gly Ala Asn Met Glu
                            -1 1
Asp Ser Val Cys Cys Arg Asp Tyr Val Ard Tyr Arg Leu Pro Leu Arg
Val Val Lys His Phe Tyr Trp Thr Ser A∮p Ser Cys Pro Arg Pro Gly
                    30
                                       35
Val Val Leu Leu Thr Phe Arg Asp Lys (lu 1le Cys Ala Asp Pro Arg
                                    50
Val Pro Trp Val Lys Met Ile Leu Asn Lys Leu Ser Gln
                            . 65
<210> 3
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<212> DNA
<213> Artificial Sequence
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<223> Description of Artificial Sequence: Primer JHSP6
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gacactatag aatagggc
                                                                  18
<210> 4
<211> 17
<212> DNA
<213> Artificial Sequence /
<223> Description of Artificial Sequence: Primer M13
<400> 4
gtaaaacgac-ggccagt
                                                                  17
<210> 5
<211> 20
<212> DNA
<213> Artificial Sequence
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<220>
<223> Description of Artificial Sequence: Primer T3.1
<400> 5
aattaaccct cactaaaggg
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<210> 6
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<220>
<223> Description of Artificial Sequence: Primer $\square{7} 7.1$
<400> 6
                                                                     22
gtaatacgac tcactatagg gc
<210> 7
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<212> DNA
<213> Artificial Sequence
<223> Description of Artificial Sequence: Primer 390-1F
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tctatctaga ggcccctacg gcgccaacat ggaag
<210> 8
<211> 33
<212> DNA
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caccggatcc tcattggctc agcttattg
                                                                     33
<210> 9
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<212> DNA
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aatggatcca cagcacggag gtgaccaag
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<223> Description of Artificial Sequence: Primer 390-3R
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<210> 11
<211> 45
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<213> Artificial Sequence
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<223> Description of Artificial Sequence: Primer 390-FX2
<400> 11
tateggatee tggtteegeg tggeecetae ggegeeaaca
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<210> 12
<211> 22
<212> DNA
<213> Artificial Sequence
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<223> Description of Artificial Sequence: Primer GEX5
<400> 12
gaaatccagc aagtatatag ca
                                                                    22
<210> 13
<211> 36
<212> DNA
<213> Artificial Sequence
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<223> Description of Artificial Sequence: Primer 390-Pel
<400> 13
attgccatgg ccggccccta cggcgccaac atggaa
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<210> 14
<211> 30
<212> DNA
<213> Artificial Sequence
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<223> Description of Artificial Sequence: Primer 390RcH
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                                                                   30
<210> 15
<211> 29
<212> DNA
<213> Artificial Sequence
<223> Description of Artificial Sequence: Primer 390RcX
<400> 15
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tggatctaga agttggcaca ggcttctgg
<210> 16
<211> 20
<212> DNA
<213> Artificial Sequence
<223> Description of Artificial Sequence, Primer DC03
<400> 16
cgaaattaat acgactcact
                                                                   20
<210> 17
<211> 67
<212> DNA
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<223> Description of Artific of Sequence: Primer
      390mycRX
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tgagaat
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<210> 18
<211> 99
<212> PRT
<213> Homo sapiens - Hu MCP-3
<220>
<400> 18
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Met Lys Ala Ser Ala Ala Leu Leu Cys Leu Leu Leu Thr Ala Ala Ala -20 -15 -10

Phe Ser Pro Gln Gly Leu Ala Gln Pro Val Gly Ile Asn Thr Ser Thr
-5 1 5

Thr Cys Cys Tyr Arg Phe Ile Asn Lys Lys Ile Pro Lys Gln Arg Leu
10 15 20 25

Glu Ser Tyr Arg Arg Thr Thr Ser Ser His Cys Pro Arg Glu Ala Val
30 35 40

Ile Phe Lys Thr Lys Leu Asp Lys Glu Ile Cys Ala Asp Pro Thr Gln
45 50 55

Lys Trp Val Gln Asp Phe Met Lys His Leu Asp Lys Lys Thr Gln Thr
60 65 70

Pro Lys Leu 75

<210> 19

<211> 99

<212> PRT

<213> Homo spapiens - Hu MCP-1

<400> 19

Met Lys Val Ser Ala Ala Leu Leu Cys Leu Leu Ile Ala Ala Thr

Phe Ile Pro Gln Gly Leu Ala Gln Pro Asp Ala Ile Asn Ala Pro Val

Thr Cys Cys Tyr Asn Phe Thr Asn Arg Lys Ile Ser Val Gln Arg Leu
10 20 25

Ala Ser Tyr Arg Arg Ile Thr Ser Ser Lys Cys Pro Lys Glu Ala Val

Ile Phe Lys Thr Ile Val Ala Lys Glu Ile Cys Ala Asp Pro Lys Gln
45 50 55

Lys Trp Val Gln Asp Ser Met Asp His Leu Asp Lys Gln Thr Gln Thr
60 / 65 70

Pro Lys Thr 75

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<211> 76

<212> PRT

<213> Homo sapiens - Hu MCP-2

80

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<220>
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·<400> 20

Gln Pro Asp Ser Val Ser Ile Pro Ile Thr Cys Cys Phe Asn Val Ile
1 5 10 15

Asn Arg Lys Ile Pro Ile Gln Arg Leu Glu Ser Tyr Thr Arg Ile 7hr
20 25 30

Asn Ile Gln Cys Pro Lys Glu Ala Val Ile Phe Lys Thr Lys Arg Gly
35 40 45

Lys Glu Val Cys Ala Asp Pro Lys Glu Arg Trp Val Arg Asp/Ser Met 50 55 60

Lys His Leu Asp Gln Ile Phe Gln Asn Leu Lys Pro 65 70 75

<210> 21

<211> 91

<212> PRT

<213> Homo sapiens - RANTES

<220>

<400> 21

Met Lys Val Ser Ala Ala Ala Leu Ala Val Ile/Leu Ile Ala Thr Ala
-20 -15 -10

Leu Cys Ala Pro Ala Ser Ala Ser Pro Tyr Ser Ser Asp Thr Thr Pro
-5 1 5

Cys Cys Phe Ala Tyr Ile Ala Arg Pro Leu Pro Arg Ala His Ile Lys
10 15 20 25

Glu Tyr Phe Tyr Thr Ser Gly Lys Cys Śer Asn Pro Ala Val Phe 30 /35 40

Val Thr Arg Lys Asn Arg Gln Val Cys Ala Asn Pro Glu Lys Lys Trp
45 55

Val Arg Glu Tyr Ile Asn Ser Leu Glu Met Ser

<210> 22

<211> '91

<212> PRT

<213> Homo sapiens - MIP-1/beta

<220>

<400> 22

Met Lys Leu Cys Val Thr Val Leu Ser Leu Leu Met Leu Val Ala Ala
-20 -15 -10

Phe Cys Ser Pro Ala Leu Ser Ala Pro Met Gly Ser Asp Pro Pro Thr
-15. 1 5

Ala Cys Cys Phe Ser Tyr Thr Arg Glu Ala Ser Ser Asn Phe Val Val 10 15 20

Asp Tyr Tyr Glu Thr Ser Ser Leu Cys Ser Gln Pro Ala Val Val Phe
30 35 40

Gln Thr Lys Arg Ser Lys Gln Val Cys Ala Asp Pro Ser/Glu Ser Trp
45 50 55

Val Gln Glu Tyr Val Tyr Asp Leu Glu Leu Asn
60 65

<210> 23

<211> 92

<212> PRT

<213> Homo sapiens - MIP-1 alpha

<220>

<400> 23

Met Gln Val Ser Thr Ala Ala Leu Ala Val Leu Leu Cys Thr Met Ala
-20 -15 / -10

Leu Cys Asn Gln Phe Ser Ala Ser Leu Ala Ala Asp Thr Pro Thr Ala
-5 1 5 10

Cys Cys Phe Ser Tyr Thr Ser Arg Gln Ile Pro Gln Asn Phe Ile Ala
15 20 25

Asp Tyr Phe Glu Thr Ser Ser Gln Cys Ser Lys Pro Gly Val Ile Phe

Leu Thr Lys Arg Ser Arg Gln Val Cys Ala Asp Pro Ser Glu Glu Trp
45 50 55

Val Gln Lys Tyr Val Ser Asp Leu Glu Leu Ser Ala
60 70

<210> 24

<211> 96

<212> PRT

<213> Homo sapiens /- I-309

<220>

<400> 24

Met Gln Ile Ile Thr Thr Ala Leu Val Cys Leu Leu Leu Ala Gly Met
-20 -15 -10

Trp Pro Glu Asp Val Asp Ser Lys Ser Met Gln Val Pro Phe Ser Arg

-5 . 1 . 5

Cys Cys Phe Ser Phe Ala Glu Gln Glu Ile Pro Leu Arg Ala Ile Leu
10 20 25

Cys Tyr Arg Asn Thr Ser Ser Ile Cys Ser Asn Glu Gly Leu Ile Phe

Lys Leu Lys Arg Gly Lys Glu Ala Cys Ala Leu Asp Thr Val Gly Trp
45 50 55

Val Gln Arg His Arg Lys Met Leu Arg His Cys Pro Ser Lys Arg Lys
60 65 70

<210> 25

<211> 93

<212> PRT

<213> Artificial Sequence - Human MDC analog

<220>

<223> The amino acid at position 24 is selected from the group consisting of arg, gly, ala, val, leu, ile, pro, ser, thr, phe, tyr, trp, aspartate, glutamate, asn, gln, cys, and met

<220>

<223> The amino acid at position 27 is independently selected from the group consisting of lys, gly, ala, val, leu, ile, pro, ser, thr, phe, tyr, trp, aspartate, glutamate, asn, gln, cys, and met

<220>

<223> The amino acid at position 20 is independently selected from the group consisting of tyr, ser, lys, arg, his, aspartate, glutamate, asn, gln, and cys

<220>

<223> The amino acid at position 50 is independently selected from the group consisting of glu, lys, arg, his, gly, and ala

<220>

<223> The amino acid at position 59 is independently selected from the group consisting of trp, ser, lys, arg, his, aspartate, glutamate, asn, gln, and cys

<220>

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<223> The amino acid at position 60 is independently
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      lys, arg, his, aspartate, glutamate, asn, gln, and
      cys
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                -20
                                                         -10
                                     -15
Val Ala Leu Gln Ala Thr Glu Ala Gly Pro Tyr Gly Ala Asn Met Glu
                                1
Asp Ser Val Cys Cys Arg Asp Tyr Val Arg Tyr Arg Leu Pro Leu Xaa
    10
                        15
Val Val Xaa His Phe Xaa Trp Thr Ser Asp Ser Cys/ Pro Arg Pro Gly
Val Val Leu Leu Thr Phe Arg Asp Lys Xaa Ile Çys Ala Asp Pro Arg
                45
                                    50
Val Pro Xaa Xaa Lys Met Ile Leu Asn Lys Leu Ser Gln
                                65
<210> 26
<211> 30
<212> DNA
<213> Artificial Sequence
<223> Description of Artifi¢ial Sequence: Primer 390-7F
<400> 26
tattggatcc gttctagctc cctgttctcg
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<210> 27
<211> 31
<212> DNA
<213> Artificial Sequence
<223> Description of Artificial Sequence: Primer 390-8R
<400> 27
ccaagaattc ctgcagccac /tttctgggct c
                                                                   31
<210> 28
<211> 20
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<212> DNA

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<213> Artificial Sequence
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<223> Description of Artificial Sequence: Primer ARA1
<400> 28
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                                                                    20
<210> 29
<21.1> 20
<212> DNA
<213> Artificial Sequence
<220>
<223> Description of Artificial Sequence: Primer ARA2
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<211> 70
<212> PRT
<213> Artificial Sequence
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<223> Description of Artificial Sequence: /Human MDC analog
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Leu Gly Pro Tyr Gly Ala Asn Met Glu Asp Ser Val Cys Cys Arg Asp
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Tyr Val Arg Tyr Arg Leu Pro Leu Arg Val Val Lys His Phe Tyr Trp
             20
                                  25
Thr Ser Asp Ser Cys Pro Arg Pro Gly/Val Val Leu Leu Thr Phe Arg
         35
Asp Lys Glu Ile Cys Ala Asp Pro Arg Val Pro Trp Val Lys Met Ile
     50
                         55
                                              60
Leu Asn Lys Leu Ser Gln
 65
<210> 31
<211> 69
<212> PRT
<213> Artificial Sequence
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<223> Description of Artificial Sequence: Human MDC analog
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15.

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Gly Pro Tyr Gly Ala Asn Met Glu Asp Ser Val Cys Cys Arg Asp Tyr
                                 . 10
Val Arg Tyr Arg Leu Pro Leu Arg Val Val Lys His Phe Tyr Trp/Thr
            20
                               25
Ser Asp Ser Cys Pro Arg Pro Gly Val Val Leu Leu Thr Phe Arg Asp
                           40
Lys Glu Ile Cys Ala Asp Pro Arg Val Pro Tyr Leu Lys Met Ile Leu
                       55
                                          60
Asn Lys Leu Ser Gln
65
<210> 32
<211> 69
<212> PRT
<213> Artificial Sequence
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Gly Pro Tyr Gly Ala Asn Met Glu Asp Ser Yal Cys Cys Arg Asp Tyr
                                  110
Val Arg Tyr Arg Leu Pro Leu Arg Val Va/ Lys Glu Tyr Phe Tyr Thr
Ser Asp Ser Cys Pro Arg Pro Gly Val /Val Leu Leu Thr Phe Arg Asp
                           40
Lys Glu Ile Cys Ala Asp Pro Arg Val Pro Trp Val Lys Met Ile Leu
                       55
Asn Lys Leu Ser Gln
65
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<400> 33
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		_				-			_			_			acc Thr 30		275
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							_						/		ctg Leu		371
								Arg		-		,			ctg Leu		419
											,				ttt Phe		467
										,					tgc Cys 110		515
•									,						ttc Phe		563
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							,								ctg Leu		707:
															aag Lys 190		755

W

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							aag Lys	_	Leu						/	803
							tta Leu 215								/	851
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							ttt Phe			,						1139
		_					gtg Val		<i>i</i> –			_				1187
							ccc									1235
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taag	gagat	tcc (ctga	gcca	gt gt	cago	gagga	a agg	gctta	acac	cca	cagt	gga (aaga	cagctt	1402
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1

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1677

<210> 34

<211> 360

<212> PRT

<213> Homo sapiens - human CCR4

<400> 34

Met Asn Pro Thr Asp Ile Ala Asp Thr Thr Leu Asp Glu Sér Ile Tyr

5 10 / 15

Ser Asn Tyr Tyr Leu Tyr Glu Ser Ile Pro Lys Pro Cys Thr Lys Glu
20 25 30

Gly Ile Lys Ala Phe Gly Glu Leu Phe Leu Pro Pro Leu Tyr Ser Leu
35 40 45

Val Phe Val Phe Gly Leu Leu Gly Asn Ser Val Val Leu Val Leu 50 55 60

Phe Lys Tyr Lys Arg Leu Arg Ser Met Thr Asp Val Tyr Leu Leu Asn 65 70 75 80

Leu Ala Ile Ser Asp Leu Leu Phe Val Phe Ser Leu Pro Phe Trp Gly
85 90 95

Tyr Tyr Ala Ala Asp Gln Trp Val Phe Gly Leu Gly Leu Cys Lys Met

100 105 / 110

Ile Ser Trp Met Tyr Leu Val Gly Phe Tyr Ser Gly Ile Phe Phe Val

Met Leu Met Ser Ile Asp Arg Tyr Leu Ala Fle Val His Ala Val Phe 130 135 140

Ser Leu Arg Ala Arg Thr Leu Thr Tyr My Val Ile Thr Ser Leu Ala 145 150 155 160

Thr Trp Ser Val Ala Val Phe Ala Ser Leu Pro Gly Phe Leu Phe Ser 165 170 175

Thr Cys Tyr Thr Glu Arg Asn His Thr Tyr Cys Lys Thr Lys Tyr Ser

Leu Asn Ser Thr Thr Trp Lys Val Leu Ser Ser Leu Glu Ile Asn Ile
195 / 200 205

Leu Gly Leu Val Ile Pro Leu Gly Ile Met Leu Phe Cys Tyr Ser Met 210 220

Ile Ile Arg Thr Leu Gln His Cys Lys Asn Glu Lys Lys Asn Lys Ala 230 235 Val Lys Met Ile Phe Ala Val Val Leu Phe Leu Gly Phe Trp /Thr 245 250 Pro Tyr Asn Ile Val Leu Phe Leu Glu Thr Leu Val Glu Leu G/u Val : 260 265 270 Leu Gln Asp Cys Thr Phe Glu Arg Tyr Leu Asp Tyr Ala Ile Gln Ala 280 285 Thr Glu Thr Leu Ala Phe Val His Cys Cys Leu Asn Pro 1/1e Ile Tyr 290 295 Phe Phe Leu Gly Glu Lys Phe Arg Lys Tyr Ile Leu G/n Leu Phe Lys 310 315 Thr Cys Arg Gly Leu Phe Val Leu Cys Gln Tyr Cys/Gly Leu Leu Gln 325 Ile Tyr Ser Ala Asp Thr Pro Ser Ser Ser Tyr Thr Gln Ser Thr Met 340 345 Asp His Asp Leu His Asp Ala Leu 355 <210> 35 <211> 1784 <2:12> DNA <213> murine MDC cDNA <220> <221> CDS <222> (1)..(276) <220> <221> mat_peptide <222> (73)..(276) <400> 35 atg tet aat etg egt gte eca/etc etg gtg get etc gte ett ett get 48 Met Ser Asn Leu Arg Val Pro Leu Leu Val Ala Leu Val Leu Leu Ala -20 -15 gtg gca att cag acc tct gat gca ggt ccc tat ggt gcc aat gtg gaa Val Ala Ile Gln Thr Ser Asp Ala Gly Pro Tyr Gly Ala Asn Val Glu - 5 - 1

gac agt atc tgc tgc cag gac tac atc cgt cac cct ctg cca tca/cgt 144 Asp Ser Ile Cys Cys Gln Asp Tyr Ile Arg His Pro Leu Pro Set Arg 192 tta gtg aag gag tto tto tgg acc tca aaa tcc tgc cgc aag/cct ggc Leu Val Lys Glu Phe Phe Trp Thr Ser Lys Ser Cys Arg Lys Pro Gly gtt gtt ttg ata acc gtc aag aac cga gat atc tgt gcc gat ccc agg Val Val Leu Ile Thr Val Lys Asn Arg Asp Ile Cys Ala/Asp Pro Arg cag gtc tgg gtg aag aag cta ctc cat aaa ctg tcc tagggaggag 286 -Gln Val Trp Val Lys Lys Leu Leu His Lys Leu Ser gacctgatga ccatgggtet ggtgtggtec agggaggete agéaagceet attettetge 346 cattccagca agageettge caacgaegee acetttacte actecatee eetgggetgt 406 cactetgtea ggetetggte cetetacete cectetatee/ettecagett atceceette 466 aatgtggcag ctgggaaaca cattcaggcc agccttaccc aatgcctact ccccactgct 526 ttagatgaga ccagcgtcct tgttttgatg ccctgatect atgatgcctt ccccateccc 586, ageettggee eeettetett ettgeatgta gggaag $m{g}$ eee ataggtttea aatatgtget 646 \sim acctacttcc ctttctgggg ggttctaata cccagcatgt ttttcctgct gcaggcacct 706 atecagtiges acacacetes caagititieta teagiteceag tyggeateca ceaageeeea 766 🚉 🔆 aacttcagac ttccttggcc tccacctact ct/cagtagaa ttctgggagt ttcaggctgg 826 tccaccaggc cccccagggt taggccaagg fccccaccagg agetectect gtttcttggt 886 ctgcagcacg gggcagggag caaggagcad gctcagaatc agattectta aaggagctgc 946 agactccatc agtaaaagga atcttt/c/c catccctgaa tataaggcag ttttctgtca 1006 acacagagac tcaggttgtt agaaatggcc acatagatca actgtgaaac cctaaattta 1066 ccaagaatca acttccaccc ctcttcáacc acatgctagg gtcttttact ttctctgccc 1126 cacacctttg actecttgee tgtgt/agetg atagtegaag ttatgetatg gtgteagtga 1186 ctgccacagt ttgtttggta ttataagcta tagttatatt tatataggaa agaggataaa 1246 tatatgtggg ccaaatagac gaactggaga gttttaggat ctgggggcag gaagggccat 1306 acaaagtgat acctcagaaa atagatggtt gtgggagctg ctgccagtgg cagagttaac 1366 ttaaagaact taattgaaat tattettgag tggetgagge caagacaaga atatagaace 1426

P 1

cattettget teectggaga caacagtggt eccaggggaa ggaataaace teettgetee 1486 teetggaggga geatggeetg rettageega gtgactggae tgtgtgagat tggggggeate 1546 getttteety teetgageete agetgacage atatgggaee acaaaggget tgatecaaac 1606 cacagggatt gacagtgeea gecacagetg tgtecaggge tegtgttetg ecagaaggag 1666 caeeetggaeg accagggeea ecactagtge taetttgete aeegeecatg eatgteetga 1726 aggteeetee eeeteete etaettetgg gaaaataaat getggeeaat aatacetg 1784

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-5 -1 1 / 5

Asp Ser Ile Cys Cys Gln Asp Tyr Ile Árg His Pro Leu Pro Ser Arg
10 15 / 20

Leu Val Lys Glu Phe Phe Trp Thr Ser Lys Ser Cys Arg Lys Pro Gly
25 30 / 35 40

Val Val Leu Ile Thr Val Lys Asn/Arg_Asp Ile Cys Ala Asp Pro Arg 45 50 55

Gln Val Trp Val Lys Lys Leu Leu His Lys Leu Ser

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	cc aat la Asn				Ser					Asp			,		96
. aat a	5 ::		cat	+ + c	10		a 2a	:.	t 2.0	15	200	+01	/	+ ca	144
	tg cca eu Pro											- /			,
	gc aag rg Lys														192
	ct gac la Asp									,					240
gcc ta Ala	agggag	aag g	ggcct	gato	ga co	cacgo	ggtci	t ggi	tgtct	cca	caag	gget:	cag		293
caaac	cctat	cctt	ctgcc	a to	ccago	caaga	a gco	cttg	ccaa	caa	ctcca	acc ·	tttg	ctcacc	353
tccat	cccct '	gggti	tgtca	ac to	ctgtg	gaago	c cto	cggg	ccc	tgta	actto	cct.	gtcc	gtcccc	413
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			1				/	//-/						tcctat	
							$\mathcal{L}/$							accaat	
						/	///							gaatat	
						1			•					gtccca	
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					/	,								ccaaca	,
,					J /									ccaatt	
tctga	anang .	aacn	gccca	at to	entec	ccntt	c aaa	aatt	aacc	ttt	cccc	ccc.	tece	tgangt	953
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11

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      5 :
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Pro Leu Pro Pro Arg Phe Val Lys Glu Phe Tyr Trp Thr Set Lys Ser
Cys Arg Lys Pro Gly Val Val Leu Ile Thr Ile Lys Asn Arg Asp Ile
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Cys Ala Asp Pro Xaa Met Leu Trp Val Lys Lys Ile Leu His Lys Leu
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Ala
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                                    -80
gca gca tcc tcc gca tta gct/gct cca gtc aac act aca aca gaa gat
Ala Ala Ser Ser Ala Leu Ala Ala Pro Val Asn Thr Thr Glu Asp
                                -65
gaa acg gca caa att ccg gct/ gaa gct gtc atc ggt tac tta gat tta
Glu Thr Ala Gln Ile Pro Ala Glu Ala Val Ile Gly Tyr Leu Asp Leu
        -55
                            -50
                                                -45
gaa ggg gat ttc gat gtt /gct gtt ttg cca ttt tcc aac agc aca aat
                                                                  194
Glu Gly Asp Phe Asp. Val/Ala Val Leu Pro Phe Ser Asn Ser Thr Asn
                       -35
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aac ggg tta Asn Gly Leu -25					Ser Ile		
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gaa gac agc Glu Asp Ser 10	Val Cys	-	_	_	,		
cgc gtg gtg Arg Val Val 25					,		
ggc gtg gtg Gly Val Val 40							
aga gtg ccc Arg Val Pro							476
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-24- . His Phe Tyr Trp Thr Ser Asp Ser Cys Pro Arg Pro Gly Va∤ Val Leu 35 Leu Thr Phe Arg Asp Lys Glu Ile Cys Ala Asp Pro Arg ∜al Pro Trp Val Lys Met Ile Leu Asn Lys Leu Ser Gln 65 <210> 41 <211> 93 <212> PRT <213> Artificial Human MDC analog <223> The amino acid at position 2 is not proline <220> <400> 41 Met Ala Arg Leu Gln Thr Ala Leu Leu Val Val Leu Val Leu Leu Ala -15/ Val Ala Leu Gln Ala Thr Glu Ala Gly Xaa Tyr Gly Ala Asn Met Glu - 5 Asp Ser Val Cys Cys Arg Asp Tyr Val Arg Tyr Arg Leu Pro Leu Arg 15 Val Val Lys His Phe Tyr Trp Thr Ser Asp Ser Cys Pro Arg Pro Gly 35 Val Val Leu Leu Thr Phe Arg Asp/Lys Glu Ile Cys Ala Asp Pro Arg 45 **5**0 Val Pro Trp Val Lys Met Ile Lew Asn Lys Leu Ser Gln 65 <210> 42 <211> 538 <212> DNA <213> Homo sapiems <220> <221> CDS <222> (53)..(334) <220>

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cca ctg aag atg Pro Leu Lys Met -20				/ /	_	106 %
cag cac atc cac Gln His Ile His -5						154
ctg gag tac ttc Leu Glu Tyr Phe 15				//	Thr Trp	202
tac cag aca tct Tyr Gln Thr Ser				/ /		250 :.
gtg cag ggc agg Val Gln Gly Arg 45			· / /			298°
aat gca gtt aaa Asn Ala Val Lys 60					tċc 🕠	344
tcaccccaga ctcct	gactg to	tcccggga	ctacctggga	cctccaccgt	tggtgttcac	404
tcaccccaga ctcct			//			
	gegeet gg	gtccaggg	gaggcettee	agggacgaag	aagagccaca	464
egeececace etgaç	gegeet gg	gtccaggg	gaggcettee	agggacgaag	aagagccaca	464
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cgccccacc ctgag gtgagggaga tccca aagtctttat cctc <210> 43 <211> 94 <212> PRT <213> Homo sapie	gegeet gg atecee tt 	gtccaggg gtctgaac	gaggcettcc	agggacgaag	aagagccaca gcccagarta Gly Ala	464 · 524
cgccccacc ctgag gtgagggaga tccca aagtctttat cctc <210> 43 <211> 94 <212> PRT <213> Homo sapie <400> 43 Met Ala Pro Leu	gegeet gg	gtccaggg gtctgaac	gaggcettee tggagceatg Leu Val Thr	agggacgaag ggcacaaagg Leu Leu Leu -10	aagagccaca gcccagarta Gly Ala	464 · 524
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20 25 30

Asp Ser Val Cys Cys Arg Asp Tyr Val Arg Tyr Arg Met Pro Leu Arg
35 40 45

Val Val Lys His Phe Tyr Trp Thr Ser Asp Ser Cys Pro Arg Pro Gly 50 55 60

Val Val Leu Leu Thr Ser Arg Asp I/ys/Glu le Cys Ala Asp Pro Arg

Val Pro Trp Val Lys Met Ile Ley Asa Lys Leu Ser Gln